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(WO/2001/005932) STRUCTURED SURFACTANT SYSTEMS

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Carbohydrates are used to confer structure on a structured Abstract:

surfactant system comprising surfactant and water and capable of suspending particles of solid, liquid and/or gas.



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(WO/2001/005932) STRUCTURED SURFACTANT SYSTEMS

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STRUCTURED SURFACTANT SYSTEMS The present invention relates to the formulation of structured surfactant suspending systems. It is particularly relevant to the formulation of personal care formulations such as shampoos and skin cleansing preparations.

STRUCTURED SURFACTANT Suspending solids in liquids presents a problem, if the solids differ in density from the liquid they will trend either to sediment or float. Increasing the viscosity of the liquid can related, but not prevent such separation, and high viscosities are generally undesirable. Colloidal systems, in which the suspended particles are sufficiently small to experience Brownian motion, e. g. liess than 1 micron, may be kinetically stable. However the difficulty or undesirability of comminuting some solids to such sizes, and the impossibility of maintaining many of them at this level in the text of crivated quowith or applicance. In the text of crivated quowith or applicance in the such contrast or provided in the such contrast or provi

Adjusting the density of one phase to match that of the other is usually impracticable.

Moreover such systems are almost always temperature-unstable due to differential rates of thermal expansion.

One method of suspension which permits even retainvely large particles to be stably suspended is shructured surfactant. The teim covers systems in which a surfactant mesophase, usually a lamelfular or G-phase, alone or more usually literapersed with an aqueous phase, provides a yield stress which is sufficient, when the system is at rest, to immobility literapersed with an aqueous phase, provides a yield stress which is sufficient, when the system to be poured like a normal liquid. Such systems any suspended particles. but which is sufficiently low to allow the system to be poured like a normal liquid. Such systems may display very low apparent viscosities when stirred, pumped or poured and yet be capable of maintening particles, sometimes of millimetre or larger size, indefinitely in suspension.

Three main types of suspending system have been employed in practice, all involving a G-phase. In which bilayers of surfactant are arranged with the hydropholic part of the molecule on the interior and the hydropholic part of the bilayer (or vice versa). The bilayers lie side by side, e. g. in a parallel or concentric configuration, sometimes separated by aqueous layers. G-phases (also Known as La phases) can usually be identified by their characteristic textures under the polarising microscope and/or by x-ray diffraction, which is often able to detect evidence of lameliar symmetry. Such evidence may comprise first, second and sometimes third order peaks with d-spacing (-where Q is the momentum transfer vector) in a simple Q integral ratio 1:2:3. Other types of symmetry by defiderent ratios, usually non-integral.

The d-spacing of the first peak in the series corresponds to the repeat spacing of the bilayer system.

Most surfactants form a G-phase either at amblent or at some higher temperature when mixed with water in certain specific proportions. However such conventional G-phases do not usually function as structured suspending systems. Useful quantities of solid render them unpourable and smaller amounts tend to sediment.

The main types of structured system used in practice are based on dispersed lamellar, spherutility and expanded lamellar phases. Dispersed lamellar phases are arranged as phase systems in which the surfactant bilayers are arranged as lameller plates to form domains of G-phases which are interspersed with an aqueous phase to form an opaque gel-like system. They are described in EP O 086 614.

Spherulitic phases comprise well defined spheroidal bodies, usually referred to in the art as spherulities, in which surfactant bilayers are arranged as concentric shells. The spherulites usually have a diameter in the range 0.1 to 15

microns and are dispersed in an aqueous phase in the manner of a classical emulsion, but interacting to form a structured system, Spherulitic systems are described in more detail in EP O 151-884.

Many structured surfactant systems are intermediate between dispersed lamellar and spherulitic, involving both types of structure. Usually systems having a more spherulitic character are preferred because they tend to have lower viscosity. A variant on the spherulitic system comprises prolate or rold shaped bodies sometimes referred to as battonettes.

Both of the foregoing systems comprise two phases. Their stability depends on the presence of sufficient dispersed phase to pack the system so that the interaction between the spherulites or other dispersed mesophase domains prevents separation. If the amount of dispersed phase is insufficient, e.g. because there is not enough surfactant or because the surfactant is too soluble in the aqueous phase to form sufficient of a mesophase, the system will undergo separation and cannot be used to suspend solidis. Such unstable systems are not considered to be obtained from the purpose of this senedication.

A hild type of structured surfactant system comprises an expanded G-phase. It differs from the other two types of structured system in being assentially a single phase, and from conventional G-phase in having a wider d-opacing. Conventional G- phases, which typically contain 60 to 75% by weight surfactant, have a d-spacing of about 4 to 7 nanometers. Alternipts to suspend solids in such phases results in stiff pastes which are either non-pourable, unstable or both. Expanded G-phases with d-spacing greater than 8, e. g. 10 to 15 nanometers, form when the electrolyte is added to acqueous surfactants at concentrations just below those required to form a normal G-phase, particularly to stratants in the M phase. The M phase comprises surfactant molecules arranged to form cylindrical rods of indefinite length, it exhibits havagonal symmetry and a distinctive texture under the polaristing microscopie.

Typical M phases have so high a viscosity that they appear to be curdy solids. M phases near the lower concentration limit (the L I/M phase boundary) may be pourable but have a very high viscosity and often a mucous-like appearance. Such a systems tend to form expanded G-phases particularly readily on addition of sufficient electrolyte. Expanded G-phases such described in more detail in E.P.O. 530 708. In the absence of suspended matter they are transucent, untilke dispersed inameliar or spherulitic phases which are necessarily opaque. They are optically anisotropic and have shear dependent viscosity. In this they differ from L, phases which are micellar solutions and which include microemulisons. L, phases are clear, coldaily isotropic and are usually substantially Newtonian. They are unstructured and cannot suspend solids.

Some L. phases exhibit small angle x-ray diffraction spectra which show evidence of hexagonal symmetry and/or exhibit shear dependent viscosity. Such phases usually have concentrations near the LLM phase boundary and may form expanded G-phases on addition of electrolyte. However in the absence of any such addition of electrolyte they lack the yield point required to provide suspending properties, and are therefore not considered to be "structured systems" for the purpose of this specification.

Expanded G phases are usually less robust than spherulihc systems. They are liable to undergo a phase change at elevated temperatures to the optically-isocropic, unstructured L2 phase. Relatively low yield stress may limit the maximum size of particle that can be stably suspended.

Most structured surfactants require the presence of electrolyte as well as surfactant and water in order to form structured systems capable of suspending sollies. However certain relatively hydrophobic surfactants such as isopropylamine alkyl benzene sulphonate can form spheruitles in water in the absence of electrolyte. Such surfactants are capable of suspending solids in the absence of electrolyte as described in EP O 414 549.

APPLICATION Structured surfactants have been applied to the problems of suspending; water insolvible or sparingly soluble huideds in is laundly detergent; antifocus and enzymes in learning detergents and other surfactant systems; abrasives in hard surface cleaners; pesticides and oils in agrochemical preparations (EP O 388 239 and EP O 488 231 cock cuttings in drilling muds (EP O 430 602), dyestuffs in dyebath concentrates and printing insik (EP O 472 989); talcs; oils and other cosmetic ingredients in personal care formulations (EP O 530 768). The present invention is applicable to accentrate the product may be a major factor in promoting sales, for example, to shampoos, body lotions, shower gels or hair creams, It may also be applied about the promoting sales, for example, to shampoos, body lotions, shower gels or hair creams. It may also be applied to the promoting sales, for example, to shampoos, body lotions, shower gels or hair creams.

concentrates for the food industry and to toothpastes.

FLOCCULATION A problem with the two phase structured surfactant systems, and especially spherulits systems, is floculation of the dispersed surfactant structures. This tends to oour at high surfactant structures of the tender of the dispersed surfactant structures. The tends to oour at high surfactant stander high electrics. It can have the effect of making the composition very viscous and/or unstable with the dispersed surfactant separating from the aqueous phase.

Certain amphiphilio polymers have been found to act as deflocculants of structured surfactants. One type of deflocculant polymer exhibits cteniform (comb-shaped) architecture with a hydrophilic backbone and hydrophobic side chains or vice versa.

A typical example is a random copolymer of acrylic acid and a fatty alkyl acrylate.

Cteniform deflocculants have been described in a large number of patents, for example WO-A-9106622,

A more effective type of defloculant has surfactant rather than centiform architecture, with a hydrophilic polymer group attached at one and to a hydrophobic group. Such defloculants are typically telomers formed by telomerising a hydrophilic monomer with a hydrophobic belogent. Examples of surfactant defloculants include alikyl thiol polyacrylates and alkyl polyglycosides. Surfactant defloculants are described in more details in EP O 623 670. In a copending PCT patent application pCT/GB000447 filled or 22 June 2000 claiming priority from British patent application, on, 9914673 we have described the use of small amounts (e. g. about 15% by weight of the composition) of carbohydrates such as sugars and alcinates as defloculants in structured surfactant compositions.

The latter comprise surfactant, water and electrolyte in proportions adapted to form flocculated two-phase structured surfactant systems in the absence of the carbohydrate.

THE PROBLEM Existing structured surfactant formulations are constrained by several limitations which have hitherto limited their application, especially in the areas of cosmetics and personal care. These include the following: -1. Unless a substantial amount of electroryte is present the choice of surfactant is limited to a fairly small range of relatively insoluble surfactants such as loopneyd alkyl benzene sulphonates. For many applications these are not the surfactants of choice from a performance point of view, and in some cases are totally inappropriate.

2. Spheruiltic or dispersed lamellar structured surfactants are opaque.

This limits the visual effects that can be achieved and may be perceived as less attractive than a clear system in some applications.

- Expanded G phases are normally opalescent, have limited suspending power and are usually formed over narrow concentrations and/or temperature ranges which make them difficult to use in practice.
- 4. At high surfactant concentrations, e. g. above 25% by weight it is difficult to make stable structured systems without using expensive defloculants and auxiliary stabilisers.

Most surfactant systems require preservatives to prevent microbial spoilage. However preservatives are expensive, ecologically undesirable, and may cause sensitivity problems for some users.

There is a need, especially in the personal care field, for a suspending system that is clear, transparent and mobile. There is a need for a system which contains high levels of surfactant but which does not require expensive delibourist. There is a need for a system that contains relatively soluble surfactants but which does not require the presence of electrolyte as a structurant. There is a need for a cleaning composition which does not require the presence of electrolyte as

THE SOLUTION We have now discovered that formulations meeting some or all of the above needs, may be obtained by

using water soluble carbohydrate to impart structure to the surfactant system, instead of or in addition to the electrolytes used hitherto. A component of the structured surfactant system which is used to impart structure to the surfactant will be refured to herein as a "structurant".

THE INVENTION Our invention provides the use of water soluble carbohydrates as structurants in structured surfactant suspending systems.

According to a second embodiment our invention provides a structured surfactant system having suspending properties which comprises a surfactant, water and a structurant characterised in that said structurant comprises a water soluble carbohydrate. The structured system may typically be a very highly expanded G- phase, a. g. one having a lameliar repeat spacing greater than 8 and usually greater than 15mm. The composition, in the absence of suspended matter, is preferably clear and fransoriert.

According to a further embodiment the invention provides a structured surfactant system having suspending properties and comprising surfactant, carbohydrate and water and having a structural feature with a repeat spacing of 20 to 50nm. The structural feature is preferably lameliar, e.g. consisting of a highly expanded G-phase.

According to a third embodiment the invention provides a composition comprising a structured surfactant system of the invention as specified above and suspended particles. The particles may be solid. liquid or gaseous and are either insoluble in the composition or present in excess of their solublity.

THE STRUCTURED SYSTEM The term'structured system's used herein means a pourable composition comprising water, surfactant, dissolved carbohydrate and any other dissolved matter, including any costructurants, which together form a mesophase, or a dispersion of a mesophase in a continuous aqueous medium, and which has the ability to immobilies usepended particles while the system is at rest, to form a pourable suspension.

The aqueous structured systems formed by the interaction of surfactants with carbohydrates include systems which are believed to be in the form of an expanded G-phase. In particular they include novel systems having a much wider repeat spacing than the typical electrolyte-structured expanded G-phases described in EP O 530 708.

The systems of the present invention comprise structures which typically show a repeat spacing between 20 and 50nm which is approximately double the repeat spacings measured for electrolyte-structured expanded G-phase, and approximately four times the typical repeat spacing in a conventional binary surfactantivater G-phase. The following discussion is based on the assumption that the structure is lamellar. We do not, however, intend to exclude the possibility that the system may comprise non-lamellar components.

Surprisingly, despite the apparent high lamellar spacing of the G-phases of the present invention, they are generally robust with good suspending power and good temperature stability. Typically the viscosity increases slightly with increasing temperature and the systems are often stable up to 70°C or higher.

The systems when fully deserated and free from suspended fine insoluble particles are generally obtainable in a substantially clear and transparent form in marked contrast to other structured surfactant systems. This can typically be achieved by vigorous centrifugal deseration and/or by gentle heating at, e. g. 60 to 80 °C.

If the amount of surfactant or of structurant is not sufficiently high, or the ratio of electrolyte to carbohydrate is too high, the structured system of the invention will be obtained as an opaque two phase system which may be spherulific or comprise dispersed 6-base or batometres.

PROPORTIONS The proportions vary depending on the nature of the surfactant and of the carbohydrate. Figure 1 shows, in schematic form, a typical phase diagram for the system coconut diethanolamide, sucrose and water. The area marked clear lamelian' represents the clear, pourable structured suspending system. The following typical proportions are expressed by weight of the total structured system 1. e. comprising the water, surfactant, structurant and any other dissolved matter but excluding any suspended solids or water-immiscible liquide.

Generally the surfactant is present in an amount of at least 2%, e. g. at least 5% especially more than 10%, by weight of the system but preferably less than 60% e. g. less than 50%, especially less than 40% more especially less than 30%. A convenient range is 3 to 25% especially 4 to 12%.

Carbohydrate structurants are usually required in substantially higher proportions than would be required for an electrolyte structurant. Preferably in the absence of electrolyte, the carbohydrate is present in a proportion of at least 25% e.g. at least 30% and usually more than 40% by weight. Concentrations greater than 65% are usually avoided. Typically the carbohydrate is less than 60%, usually less than 55% by weight of the composition. When electrolyte is present the carbohydrate may be present in substantially lower concentrations as a costructurant. Such systems may be scherulitic but in the presence of more than about 10% sugar, do not tend to flocculate. The systems of the invention require the presence of the carbohydrate in order to form a structured suspending system. Typically the less soluble the surfactant. the less carbohydrate is required.

The proportion of water is usually greater than 20% by weight, more commonly greater than 30%, typically greater than 40% of the system, but is preferably less than 65% usually less than 60%, e. q. less than 55%.

One way of preparing suspending systems according to the invention is to prepare a G or M phase agueous surfactant and add sugar until the system clears. The G or M phases are located using conventional means, as described for example in GB2013235.

Suspending power may be quickly checked by shaking air into the sample and noting whether the bubbles remain suspended. Confirmation that the system is a true structured system and not merely a slowly separating system may be obtained by allowing the sample to stand overnight at 50 or 60 °C. If the dispersed phase has not separated out in that time, the system may be assumed to be structured. It is generally found that mixtures of two G-phase systems according to the invention also form G- phases according to the invention.

If in any case difficulty is encountered locating a sugar-structured phase according to the invention, it is usually possible for resolve by adding a minor proportion based on the weight of sugar of a co-structurant has discussed below.

THE CARBOHYDRATE The preferred carbohydrates are mono and disaccharide sugars such as sucrose, glucose or fructose. Other sugars which can be used include mannose, ribose, galactose, allose, talose, gulose, idose, arabinose, xylose, ivxose, erythrose, threose, acrose, rhamose and cellobiose. The carbohydrate may be a th-or tetra-saccharide or a water soluble polysaccharide such as soluble starch. The term carbohydrate as used here includes water soluble nonsurfactant derivatives of carbohydrates such as carboxylic acids and their salts, e. g. gluconic acid, mannic acid, ascorbio acid and aldinates or reduced sugars such as sorbitol, mannitol or inositol. The levels of carbohydrate are preferably sufficiently high to inhibit microbiological growth in the medium and preferably sufficient to act as an effective blodegradable, non-allergenic preservative for the composition, thereby obviating the need for less environmentally friendly additives.

CO-STRUCTURANT Some surfactants, especially the more water soluble surfactants such as alkyl ether sulphates form the clear lamellar phase more readily in the presence of a co-structurant. The co-structurant is preferably an electrolyte. Any water soluble salt which tends to lower the solubility of surfactant in water may be used, such as sodium tripolyphosphate, sodium carbonate, sodium clirate, sodium chloride or the corresponding potassium or ammonium salts. Alkalis such as sodium or potassium hydroxide may also be used. Other structurants include polar water-immiscible solvents such as phenolethoxy ether or a terpene, water soluble mono and dihydroxy alcohols and ether alcohols such as glycerol, propylene glycol, ethylene glycol monomethyl ether and diethylene glycol monomethyl ether.

The constructurant, if required, may in principal, be present in concentrations up to 30%, but is preferably less than 20% e. g. 0.1 to 15% by weight. Often traces of costructurants e. g. 0.1 to 3%, typically 0.5 to 2.5% by weight based on the system are sufficient, although higher concentrations can be present. For example in some perfumed systems, the solvent in the perfume may be sufficient to provide any desired co-structuring effect. We prefer that the proportion of costructurant be less than the proportion of carbohydrate, preferably less than half the proportion of carbohydrate, e. q. less than one quarter the proportion of carbohydrate. Large amounts of electrolyte are generally undesirable because they inhibit the formation of clear phases.

Generally the costructurant is present in proportions insufficient to form a stable structured system in the absence of the carbohydrate.

THE SURFACTANT The surfactant preferably comprises non-onlo surfactants such as C8 25 alkyl mono or dietheranolamides or 1 to 50 mole ethoxylates such as C8-25 alcohol of rafty acid ethoxylates, alkyl amine ethoxylates, or glyceryl or sorbitan ester ethoxylates, or polyoxypropylene oxyethylene block copolymers. Ethoxylates typloally contain from 2 to 4 6e, 3 to 30 especially 6 to 15 oxyethylene groupe. Other non-ionic surfactants include alkyl polycodies, sugar esters or amine oxides. The non-ionic surfactants typically have a HLB of from 5 to 16, e. g. 6 to 15, especially 8 to 14, e. g. 10 to 12.

However surfactants with HLB as low as 1 may be used.

The surfactant may optionally be or comprise an anionio surfactant such as an ether sulphate, an alkyl benzene sulphonate, an alkyl sulphate, alkame sulphonate, olefin sulphonate, sulphosuccinate, sulphosuccinatens, coap, sarcosinate, Itauride, isettionate, alkyl phosphate, or alkyl ether carboxylate. In each case the surfactant comprises an 8 to 25 cerbon alkyl group or alkenvil group or polypropyleneoxy group.

Alkyl or alkenyl groups may be straight or branched chain, primary or secondary and preferably have from 10 to 20 eg. 12 to 14 carbon atoms. Ether groups may comprise glycenyl groups and/or 1 to 20 mol polyoxyethylene groups e. g. 2 to 10 mole. The anionic group is usually a suiphate or sulphonate group, but may also be for example, a phosphate, phosphonate or carboxylate group. The counter ion of the anionic surfactant is usually socilium but may also be potassium, lithium, ammonium or, calcium or other alkali metal or alkalitine earth metal.

The surfactant may be or may comprise an amphoteric surfactant such as betaine, sulphobetaine or phosphobetaine, Examples include fatty alkyl dimethyl betaines, alkyl amidopropyl betaines and immidazoline betaines.

The surfactant may, alternatively be or comprise a cationic surfactant such as a Cg-25 straight or branched alkyl or alkenyl or alkylphenyl tri Ci 4 alkyl or hydroxyalkyl ammonium salt, or di Ci 4 alkyl benzyl ammonium salt, or an Cs 20. alkyl or alkenyl amide amine.

ELECTROLYTE The presence of electrolyte is not normally required for structuring but is generally blerated if required for other purposes. We particularly prefer electrolyte-free or low electrolyte, (e.g., 1 to 5% by weight) composition presonal care applications or where clear formulations are required but can tolerate much higher levels, e.g. up to 20% or more if required. For example industrial cleaning formulations may require high levels of alkali such as sodium hydroxide, carionate or silicate. The presence of builders such as clirate, potassium pyrophosphate, or sodium tripolyphosphate may also be tolerated. Electrolyte may contribute to the structuring of the composition, and may be desirable as a costructurant when very water soluble surfactants or sufficients of high HLB are used.

SUSPENDED MATTER The composition may contain suspended solid, liquid or gaseous particles. For instance the composition may contain suspended oil droip lets. The oil is preferably a mineral oil (e.g. a low molecular weight petroleum oil) or a fatty glyceride or other ester such as lauryl acetate, a terpene oil such as timonene or a silicone oil. Mixtures of oils may be used, Particularly preferred are vegetable cils such as occordul, evening primose, groundnut, tradeow foam, apricot kernel, peach kernel, avocado, jojoba and oilve oil. Oil solubie cosmetio or topical pharmaceutical ingredients may be dissolved in the oil including antiseptics, styptics, antidandruff agents such as a zino omadine (zino pyritinone) and selenium disulphide, profeins, emoillients such as lancoint, isopropyl myristate, glycenyl isocierate or propylene glycol distearate, dyss, perfures and waxes. Water inscluble particulate solids may be suspended including eriolatis such as talc, clays, polymer beads, sawdust, silica, seeds, ground nutshells and calcium phosphate, pearliers such as miss or glycenol or ethylene glycol di-stearate, glyter diditives and sunscreens such as titanium dioxide. Porous particles (so called micro-sponges) containing absorbed active ingredients or gelatin or other microcapsules may also be suspended. Other active ingredients which may be suspended including eriolation and lorgical pharmaceutical preparations, o greparations for treatment of acne, fungicides for athlete's foot or ringworm or antiseptics or antihistamines. Pigments, such as the incon oxides, may also be added.

The structured suspending systems of the invention may be used to suspend builders such as zeolite or sodium

It polyphosphate, agricultural and horticultural pestioides, blooides for water treatment, cuttings or shale in drilling music, antiforans, explosives gumes such as gum benocin, quan eacade, gum tragacenth vanthan and guar gum, enzymes, flavouring and vitamin concentrates, calcium phosphate for toothpaste, pharmaceuticals, and machinery and outling abrasives such as emery or diamond powder.

The composition may contain liquefied propellant gas dispersed in order to provide foams such as shaving foam, on release from a pressurised pack.

PEARLISING The compositions of the invention are particularly useful for suspending pearlising agents. Pearlisers are required as concentrates for incorporation into liquid formulations such as sharipoos and toiletries to import a nacrecus indescence which is attractive to consumers, and can mask inhomogeneities in the formulations.

Pearlisers typically comprise small, thin, transparent platelet crystals which can be suspended in a parallet configuration. When so suspended light failing on the crystals undergoes complex multiple reflections within the substrate similar to those which occur in a pearl and giving rise to similar optical interference effects.

Natural poarls comprise alternate layers of calcium carbonate and protein. Artiflicial pearlisers which can be suspended according to the invention include guanine/hypoxenthine crystals extracted from fish scales, mea, various stats of lead, zinc, mercury and bismuth (e.g. bismuth expectively), titanium oxide and various fally acid derivatives such as magnesium stearate, occount monoethenolemice, ethylene glycol distearate and ethylene glycol monostearate. Fish scale extracts are too expensive and the inorganic pearlisers are either too toxic for general use in tolietrise e.g., etad, mercury, or relatively ineffective e.g., bismuth. The fatty acid derivatives are therefore now the most widely used pearlisers. In addition to the chemical nature and physical form of the pearliser the manner in which it is suspended has an important effect on its visual impact.

Difficulty is sometimes encountered obtaining the desired effect when incorporating pearlisers into aqueous formulations.

Conventional fatty acid derived pearlisers are supplied as solids which are usually added to a heated formulation above their meiting point and recyclatilised in eitu. The conditions of crystallisation and especially the amount and nature of the agitation applied must be carefully controlled in order to obtain an acceptable result. This makes it difficult to obtain consistent effects and renders solid pearlisers inconvenient to use.

Attempts have heen made to prepare liquid concentrates or suspensions which can be addled directly to shampoo formulations without heating. While more convenient for the user, such concentrates face the manufactures with problems of obtaining a high and consistent pearl effect, similar to those which confront the user of convenience to pearliesrs. Difficulty is also encounteded in maintaining the pentilese in stable suspension and preventing socientation.

We have now discovered that carbohydrate structured phases of the invention have the capacity to form stable suspensions of pearlisers.

The peailser may be dispersed in the equeous structure surfactant system e.g. by gently stirring, but in the case of the tatty acid derivatives are preferably prepared in situ by healing above their metiliting point. e.g. temperatures between 65 and 80%, dispersing the liquid pearliser in the structured surfactant system, preferably with sufficient stirring to form displays of the model of the preferably with sufficient solding is relatively stow e.g. the mixture is allowed to cool naturally. The amount of pearliser can be varied considerably, the main constraint on the upper limit being the viscosity.

The amount of pearliser should not be so high as to render the product unpourable, or unacceptably viscous. We prefer on economic grounds that the pearliser is present in amounts greater than suspending surfactant. Generally pearliser may be present in amounts ranging from 5% up to about 50% e.g. 10 to 45% of the total weight of the mixture.

OTHER INGREDIENTS The composition may contain minor amounts of other ingredients such as dyes, perfumes, soil suspending agents or optical brighteners. Solvents such as ethanol or isopropyl alcohol ethylene glycol, isopropylene

glycot, glycerol or water miscible glycol eithers such as ethylene glycol monomethyl either, diethylene glycol monomethyl either, diethylene glycol, and hydrotypes such as GF6 G8 flay! benzene suphponates or usen avap he required for special applications, e.g. as perfume enhancers but if not so required are generally undesirable and are preferably absent but may be tolerated in small amounts, preferably less than 10%, e.g. less than 5%, most preferably less than 10%, e.g. less than 5%, most preferably less than 10%.

OPTICAL VARIEGATION The present invention is particularly adapted to providing optically variegated fluid surfactant compositions.

Many fluid surfactant-containing products are purchased by the consumer on the basis of factors which include the appearance of the product. Detergents, shampoos, tolletries, soaps and other surfactant-based consumer products often depend upon appearance and peckaging for at least part of their consumer appeal. Striped foothpaste, marrhold soap and blue speckled detergent powder are well known examples of products whose successful promotion was based on a characteristically varienated an appearance.

However it is not, on the face of it, possible to produce any kind of lasting variegation in an otherwise homogenous, pourable, liquid formulation.

It is known to combine two or more immiscible liquids of different density and colour to form a product which segregates into horizontal bands.

The visual effect achievable by this method is limited and the product has the practical disadvantage that the essential functional components are not evenly distributed between the different bands so that the product performs inconsistently if its not vigorously adulted immediately prior to use.

Structured surfactants may be used to suspend coloured granules, to produce a speckled effect. We have now further discovered that where two or more portions of a structured surfactant such as those according to the present invention are separately coloured by including in at least one of said proportions a pigment, which is insoluble in said continuous phase or a dye which is insoluble in said continuous phase and soluble in or absorbable on any dispersed phase and said proportions are charged to a transparent container in such a way as to produce a variegated appearance, little or no migration of pigment or dye through the composition is observed in the undisturbed sample even after prolonged standing. The variegation thus remains stable to a remarkable degree. Furthermore, provided the container is substantially full, even the agitation coused by normal handling during distribution does not significantly affect the variegated appearance of the product. Yet the product may be, to all appearances, a thin, mobile liquid.

According to a further embodiment our invention therefore provides a packaged fluid surfactant-containing product, comprising an at least partially transparent container, and therein a stable structured surfactant comprising a continuous phase and a dispersed phase and having a variegated appearance caused by the inclusion in localised portions of said structured surfactant of a dye or pigment which is insoluble in said continuous phase and present (A) as particles suspended in said continuous phase and having a particle size sufficiently small to be able to give said portions a substantially homogeneous appearance which is visually distinct form other portions of said structured surfactant in said container, and/or (S) dissolved in or absortioned on said dispersed phase.

The effects obtainable can be extremely varied. Depending upon how the different visually distinct portions of the structured surfactant are charged to the container it is possible to obtain horizontal or vertical stripes, vertical segments, marbling, bands, whorts or numerous other decorative effects.

Any pattern or opicial effect which can be instantaneously obtained by charging visually distinct liquids to a transparent container can be rendered substantially permanent, at least until the product is poured from the container, busing the structured surfactants as aaid liquids. With suitable filling techniques, it is even possible to produce readable characters so that liquid roducts may be marked with Trademarks, logoe or similar devices.

The pigment or dye may for example be, or comprise a water insoluble pigment, having a particle size preferably less than 150 microns especially less than 100 microns, most preferably less than 50 microns e.g. 0.1 to 20 microns.

The proportion of pigment required is generally small. Any pigmented portion normally only requires from 0.001 to 1% by weight of pigment to produce a sufficient effect e. g. 0.01 to 0.5% more usually 0.02 to 0.1%. The precise amount will depend on the choice of pigment and the intensity of oliour required.

The pigment may be a white pigment (e. g. titanium dioxide) a black pigment (e. g. carbon black), coloured pigment such as any water insoluble pigment hitherto used in cosmetics or detergents, or a pearlising agent such as mica.

Typically the structured surfactant has an aqueous continuous phase and the pigment or dye is water insoluble. Water insoluble dyes typically dissolve in or are absorbed on a dispersed surfactant phase.

We prefer that, apart from the playment or dye, the differently coloured portions of the structured surfactor product of our invention should have essentially the same composition. This assists maintaining the stability of the product and ensures uniform performance. However it is possible e.g. where substantially water insoluble active ingredients uniform that product or dissolved or diseased or offensive contained in a suspended oil phase, to include such ingredients in only some portions of the product. This may be useful in segregating mutually incompatible components and may permit promotional clients that the policyred portions are associated with a specific beneficial effect.

The method of filling the container determines the effects produced. For example horizontal stripes can be obtained by running a single bladed strine gently in a container partly fillied with one coloured structured surfactant and hijecting a contrasting colour through a syringe at the level of the striner. Progressively raising the striner and repeating the process produces a putrality of horizontal stribes.

Injecting the contrasting liquid while drawing the syrings up the side of the container produces a vertical stripe. Inserting a partition into the container, and filling different portions on either side, before withdrawing the partition, produces contrasting vertical halves. A multiple partition permits vertical segments. Alternatively vertical effects may be produced by inserting two or more tubes into the container and gradually withdrawing the tubes while charging the container with different coloured portions, each at the same rate.

Partially stirring two contrasting portions together before filling the container gives an attractive marbled effect. The foregoing are merely an indication of some of the different filling techniques and associated visual effects possible according to the invention. Numerous other possibilities will be apparent to those skilled in the art.

The container may be any jar, bottle, tube, sachet or other conventional container for surfactant based products. It may lypically be of glass or plastic or other transparent material, it may be oblived but is preferably at least partly clear to enable the decorative contents to be easily seen. It is possible to use deformable containers usual as explezes tubes or sachets, provided that they are sufficiently filled to give them a degree of rigidity enough to avoid loss of variegation on normal handling prort to use, but it is preferred to use rigid or at least substantially non-deformable materials.

The invention will be illustrated by the following examples in which all proportions are based on weight percentages of active ingredient based on the total weight of the composition, unless stated to the contrary. Examples 1 to 4 The following formulations were prepared: -1 2 4 C12 14 3 illoiesthersulphate 10 12.5 15 17.5 Sucrose46464646 frisodiumchrate 2 2 2 Perfume6556 Waterbalancebalancebalance All ricor formulations were clear or slightly hazy, mobile structured liquids with good suspending properties. Suspensions of tals, milheral oil, pigment, small beads and plastic novelty items were prepared. All were stable after proloned storage.

Each of the examples 1 to 4 was re-prepared (a) without the perfume (b) without the dirate and (c) without perfume or ortate. No suspending power was exhibited by any of the eight samples of preparations (a) and (c). The samples of preparation (b) all exhibited similar suspending power to the original examples. On heating the samples to 70 °C and subsequent occling a clear transparent composition was obtained.

A clear sample of Example (th) and a sample containing a red pigment in suspension were slowly poured hin a sample jar in a series of alternating additions. The refer was to protoce a sequence to horizontal stripe. When the hot was jar in a series of alternating additions. The refer was to protoce a sequence to horizontal stripe. When the hot was filter than the stripes retained their integrity and showed no signs of blurring or diffusion after one year storage including intermittent jeroidos of cerelle bakknar and six months weeks stored on its side. Example 5 A schematic phase diagram was prepared for the system coconul diethanelamide/sucrose/wate and is reproduced as fill of the drawings. The area marked/clear lamellar/represents transparent expanded G-phases havin a suspending power according to the invention, and the area marked/lamellar/comprises opaque expanded lamellar suspending systems according to the invention.

The phase boundaries illustrated were not all precisely determined.

Example 6 A sample was prepared comprising 10% excount diethanolamide, 35% water and 55% sucrose. The product was a clear attenuated G-phase with good suspending power.

Example 7 A composition comprising 10% Cl2 l4 alkyl 2 mole ethoxysulphate, 33% water, 50% sucrose, 5% ethanol based perfume, 2% sodium citrate gave a clear, attenuated G-phase with good suspending powers.

Example 6 (Shampoo formulation) 4 parts by weight of the composition of Example 7 and 1 part of the composition of Example 6 were mixed together to form a clear composition of the invention with good suspending powers and good performance as a skin cleaner and shampoo.

Example 9 (laundry datergent formulation) % Active Ingredient by weight CI2 I4 linear alkyl benzene sulphonate 6,6 Triethanolamin launyl sulphate 1,65 CI2 I4 alkyl 3 mole ethoxylate 1,6 Sucrose 55.0 Sodium diethienetiamine pentakis (methylenephosphonate) 0.55 Water balance The product was a hazy, readilly pourable liquid with good suspending power.

Example 10 (Poarl concentrate) A pearl concentrate was obtained by heating a formulation comprising 54% by weight sucrose, 10% by weight coconut di-ethanolamide, 10% by weight ethylene glycol distearate and 26% by weight water to 70°C and cooling.

A spontaneously pearly suspension was obtained.

Example 11 10% by weight Cl2 alkyl six mole ethoxylate (HLB = 10), 54% by weight sucrose and 36% by weight water, were mixed and warmed to 70 °C.

The cooled product was a clear transparent, pourable system with good suspending properties.

Example 12 % w/w succrose 40.0 perfume 2.0 sodium C) 2-14 alkyl 3 mole ethoxy sulphate (70%) 8.0 coconut dilethanolamide 2.0 sodium chloride 6.0 water balance The above formulation provided a clear, transparent, pourable fluid. Whorls of three different coloured pigments were introduced into this formulation with a synilation with a synilar way.

After three months no diffusion of the pigment was observable.

Example 13 % vivis succrose 4.0.0 perfume 2.0 trisodium oltrate dihydrate 2.0 codium CI214 alkyl 3 mole ethoxy suiphate (70% by wt active) 8.8 glycamate 5.06 occorut monoethanolamide 1.1 occorut amido propyl betaline 4.18 culum C8.0 alkyl polyglycosiole dp.1.6 (65%) 6.2 codium othoride 0.66 sodium ethylene diamine letracetate 0.05 water balance The composition was a pourable, clear, transparent fluid with suspending properties.

A plurality of coloured, polystynene beads (limit diametre) were dispersed in the composition. The suspension remained stable after three months.

Example 14 % w/w sodium CI2.4 alkył (3 mole elhoxy sulphate) 10.0 coconut diethanolamide 2.5 fructose 50.0 water balance. The above composition was a clear isotropic LI micellar solution which was unsaturated and had no suspending power. Addition of 6% by weight sodium chioride, gave a stable, easily pourable fluid, composition which after shaking was capable of suspending air bubbles. The aerated composition was stood overnight at 50°C. The aged composition was clear and transparent and maintained the air bubbles in a stable suspension. Equivalent compositions with 2 and 4%

respectively of sodium chloride were not able to suspend bubbles under the foregoing conditions.

Example 15 % w/w sodium Cl2 I4 alkyl-3 mole ethoxy sulphate 8.0 A. I. coconut diethanolamide + 10% by wt glycerol 2.0 A. I. fructose 40.0 water balance The above composition was not capable of maintaining particles in suspension.

Addition of incremental amounts of sodium chloride gave the following results: wt % sodium chloride 2 LI phase bubbles rise 4 LI phase bubbles rise 6 clear suspending phase. Bubbles suspended after ageing at 50 °C overnight.



IP SERVICES



Notices

Documents

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Description

Search result: 4 of 4

(WO/2001/005932) STRUCTURED SURFACTANT SYSTEMS Claims

Note: OCR Text

National Phase

CLAIMS 1. The use of water soluble surfactants as structurants in structured surfactant systems.

- 2. A structured surfactant system having suspending properties which comprises a surfactant, water and a structurant characterised in that said structurant comprises a water soluble carbohydrate.
- 3. A system according to claim 2 which consists of a G-phase.
- A system according to claim 3 wherein said G-phase has a lamellar repeat spacing greater than 15nm.
- A composition according to claim 4 which is transparent.
- 6. A structured surfactant system having suspending properties and surfactant, carbohydrate and water and having a structural feature with a repeat spacing of from 20 to 50nm.
- 7. A system according to claim 4 wherein said feature is lamellar.
- 8. A system according to any of claims 2 to 7 wherein said carbohydrate is present in any amount of at least 25% by weight of the mixture of water surfactant carbohydrate.
- 9. A system according to any of claims 2 to 8 wherein the surfactant is present in an amount of from I to 60% by weight of the system.
- 10. A system according to any of claims 2 to containing more than 20% by weight water.
- 1 I. A system according to claim 10 containing 30 to 60% by weight water.
- 12. A composition according to any of claims 2 to 11 wherein the carbohydrate is a mono or disaccharide sugar, gluconic acid, mannic acid, ascorbic acid, sorbitol, mannitol or inositol.
- 13. An aqueous suspension comprising a system according to any of claims 2 to 7 and particles of solid, liquid or gas stably suspended therein.
- 14. A composition according to claim 13 wherein said particles comprise a builder.
- 15. A composition according to either of claims 13 and 14 wherein said particles comprise an abrasive.
- 16. A composition according to any of claims 13 to 16 wherein said particles comprise a pesticide.
- 17. A composition according to any of claims 13 to 16 wherein said particles comprise an oil.

- 18, A composition according to any of claims 13 to 17 wherein said particles comprise a pigment.
- 19. A composition according to claim 18 having a plurality of differently pigmented zones producing a variegated visual effect.
- 20. A packaged fluid surfactant containing product, comprising an at least partially transparent container, and therein a stable structured surfactant comprising a continuous phase and a dispersed phase and having a variegated paperance caused by the inclusion in localise portions of said structured surfactant of a dye or pigment which is insoluble in said continuous phase and present (A) as particles suspended in said continuous phase and present sufficiently small to be able to give said portions a substantially homogeneous appearance which is visually distinct form other portions of said structured surfactant in said containuous, and/or (B) dissolved in or absorbed on said dispersed phase.